

EC Declaration of Conformity

Manufacturer:

Name: HANGZHOU ALLTEST BIOTECH CO., LTD.

Address : #550, Yinhai Street, Hangzhou Economic & Technological Development Area, Hangzhou -310018, P.R. China

European Representative: Name: MedNet EC-REP GmbH Address: Borkstrasse 10, 48163 Muenster, Germany

Product Name: COVID-19 Antigen Test Cassette Cat.No.: FI-NCP-502 Analyte: SARS-CoV-2 Nucleocapsid protein antigens in human Nasopharyngeal Swab Analyzer/Reader: Fluorescence Immunoassay Analyzer Model: Cassette Classification: Other Device, non-listed in Annex II of IVDD 98/79/EC Conformity Assessment Route: IVDD 98/79/EC Annex III (Excluding point 6) EDMA Code: 15 70 04 01 00

We, HANGZHOU ALLTEST BIOTECH CO., LTD., herewith declare that we are exclusively responsible for this declaration of conformity. We herewith declare that the above mentioned products meet the transposition into national law, the provisions of the following EC Council Directives and Standards. All supporting documentations are retained under the premises of the manufacturer.

DIRECTIVES

General applicable directives: DIRECTIVE 98/79/EC OF THE EUROPEAN PARLIAMENT AND OF THE COUNCIL of 27 October 1998 on in vitro diagnostic medical devices

Standard Applied: EN ISO13485:2016, EN ISO14971:2019, EN 13975:2003, EN ISO 18113-1:2011, EN ISO 18113-2:2011, EN 13612:2002/AC:2002, ISO 17511:2020, EN ISO23640:2015, EN 13641:2002, EN ISO 15223-1.2021

Place, Date of Issue: in <u>Hangzhou</u> on 09/12/2020 Date of Issue of DOC: <u>22/02/2022</u>

Signature: Name: GAO FEI (Position: General Manager)

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